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- Polyorganosiloxane based interpenetrating network polymers and methods of making.
- ② Cross-inked polyorganosiloxane networks are prepared by dispersing vinylsiloxane and silicon hydride components in a liquid monomer, cross-linking his elioxane components by hydrosilystilox, and polymerating of the organic monomer. Poly-penetraling network systems may be prepared by cross-linking one or more organic polymers is the polyorganosiloxene network. At least one of the organic polymers is a vinyl or opportune which may be polymerated by free radical polymerization. The interpenetraling polymer retwork systems may be cast, motified, extruded or formed into films, membranes and coatings. The resulting products have improved mechanical, gas permeability and optical properties compared to conventional silicone elastiments.

EP 0 315 836 A2

POLYORGANOSILOXANE BASED INTERPENETRATING NETWORK POLYMERS AND METHODS OF MAK-

### Field of the Invention

The present invention relates to the preparation of novel organosilicone interpenetrating polymer systems. More particularly, the invention is directuct to highly relaxed and perturbed polyorganosiloxanes which are cross-linked to form networks which hieropenetrate another polymer or polymers.

# Background of the Invention

Traditional silicone rubber technology has to go through the following steps: (1) polymertzation of a silicone; (2) filter compounding; and (3) transferring this compound to a mold which is maintained at high 16 temperatures for extensive periods of time under pressure. Due to the high degree of cross-linking needed to achieve useful properties, curing at elevated temperatures over an extended period of time is required. This processing is extremely slow and labor intensive. Furthermore, due to the inorganic filler and multi-functional crosslinks needed, a rubber product with low recovery and high permanent set mey result, particularly find proprepting post cured.

Previous work of Kiempner et al., Speriing, and Lipatov et al. in the 1970s on interpenetreting polymer network materials dealt with heterogeneous systems prepared either by latex blending, mell octusion or post-swelling of cross-linked polymer. Such techniques result in heterogeneous cross-linked systems with large macro domains of each of the phases present, as well as strained networks dispersed within the meterial. Due to the heterogeneous morphology present in those systems described in the literature, such envolves have been considered as blends of polymer/polymer composite systems rather than true molecularly interpenetrating networks, which were first defined by J. R. Millar, J. Chem. Soc., 1311 (1980). However, such networks depend upon re-swelling previously formed networks which results in streiped elastometic systems having over treatability.

Blends of silicone rubber and an organic vinyl polymer have also been reported, but the poor omechanical properties of subt blends due to the incompatibity of the polymeric components, resulted in no useful materials. A more recent approach is described in U.S. Patent 4.500,688 of Arkles, assigned to the same assignee es the present invention. In that patent polysitoxanes are vulcanized cross-linked or chain extended) after mixing with a matrix polymer and during met processing of the composition to produce silicone semi-interpenentating networks in the thermopleatior matrix. However, the earlier at produces a macro cell structure as described by Sperling in Macromodecula Reviews, 12:141 (1977) and interpenentating Polymer Networks and Related Materials, Plenum Press, NY, (1981). In general, the larger the domains, the poorer will be the physical properties due to the lack of penetration through the domains.

Attempts to prepare cross-linked composites of polydimethylsiloxane with polystyrane and polymethacrylste by cross-linking polysiloxane unbber, followed by swelling and polymerizing the second-40 component during high temperature peroxide curing, resulted in multiphase, densely cross-linked systems where the overall system is cured together, forming a brittle, semi-rigid or learney material with poor elastic properties. See S.L. Aggarwal, Ed., Block Coploymers, Plenum Press, NY. (1970), Such cured composites have no practical usage due to poor proporties. Earlier work does not teach the art of preparing a welldefined, perturbed and relaxed network of polyorganosiloxane as a route to preparing true interpenetrating 3 networks on a molecular scale.

### Brief Summary of the Invention

According to the invention, compositions are provided in which a relaxed, perturbed, cross-linked polyreganositosene network is dispersed in a second organic polymer, perfearby a homogeneous virily virilytidene polymer, to form a penetrating network in the second polymer. The compositions may be monopenetrating networks, where the second polymer is not significantly cross-linked; dual-penetrating networks. where the second polymer is substantially cross-linked to itself, or poly-penetrating polymer networks, where two or more polymers in addition to the polyorganosiloxane are present, with each polymer being substantially cross-linked to the cinsel flux not significantly cross-linked to the other polymers.

Compositions of the invention are produced by dispersing a component having silicon hydride (SIH) groups and an organositosone having virily groups in a reactive fleuid monomer, reacting the silicon hydride and viryl groups by hydrosilystation to form the organositoxane network in the monomer, and simultaneously or subsequently polymersizing the monomer. Dual and poly- penetrating networks are formed by sequentially cross-thinking the second polymer and any additional polymers which are present in the composition. The hydrosilystation reaction is preferably catalyzed by a pilatinum catalyst, and the second and any additional polymers may be polymerized and cross-linked by the readical, anniori, cationic or organometallic polymerization. Further, the reactions of the present invention may be carried out by modified bulk, solution or suspension polymerization polymerization.

The compositions of the present invention may be produced in the form of a pourable gel for preparing films, costings and castings, or dried beads or pellets for modding and extunion. Homogeneous compositions according to the invention may be prepared to produce optically clear compositions for contact lenses and other medical and optical uses.

# Detailed Description of the Preferred Embodiments

The present invention is directed to the preparation of inter-penetrating network systems composed of an organositionane polymer and at least one other organo polymer to produce highly retractable interpenetrating silicone disastemer systems with improved elastiomeric properties. According to rubber elasticity as theory, cross-finked, strained networks undergo less deformation and have greater retractive stress than relaxed networks. In contrast, the present Invention produces organosilicone elastiometric interpenetrating networks having significantly better recovery than both silicone rubbers and other interpenetrating network polymers prepared by the prior art.

While applicant does not wish to be bound by any particular theory, the following is believed to be at least a partial explanation for the superior properties of the compositions of the present invention. In the present invention, the interpenetrating networks are produced without swelling or shear stress being present, resulting in relaxed interpenetrating polymer networks. In the method of U.S. patent 4;800,889 of Arkies, referred to above, the vulcanization of the hydride-containing silicons via a hydrosilystion reaction takes place within a polymeric thermoplastic matrix. Although the hydrosilystion reaction in such systems is a carried out while the matrix polymer is in a fluid melt state, it is believed that a certain amount of strain is introduced into the resulting silicone interpenetrating networks due to the polymeric nature of the thermo-

In contrast, according to the present invention, the hydrosilytation reaction is carried out with the vinycontaining organosiloxane and the silicon hydride component being dispersed, and preferably dissolved, in 40 a monomer for the second polymer. This hydrosilytation reaction is carried out, for example, by using platform catalyst systems such as those described in U.S. patent 3,419,893. This cross-linking of the vinytistloxane and silicon hydride components is believed to result in a substantially relaxed, perturbed polygragnosiloxane network which will penetrate or interperetrate subsequently formed polymers from the monomer(s) in which the organosiloxane components are dispersed.

48 used herein, the term "perturbed" refers to the extended state of the polyorganosiloxane chains as allowed by the fluid medium of the fliquid monomer in which the organosiloxane components are dispersed and cross-linked. This contrasts with an unperturbed polymer state, such as obtained when a polymer is formed or cross-linked in a conventional bulk polymerization, whereby the polymer is in a substantially coiled or non-extended state. The concept of perturbation is discussed mathematically, for example. In PJ. 59 Flory, Principles of Polymer per 176, Esseiver, Amsterdam (1978). Amsterdam (1978) or Polymers, page 176, Esseiver, Amsterdam (1978).

Similarly, as used herein, the term "relaxed" refers to the substantial absence of stress during the cross-initing of the organosioxane components so that the resulting polyogranosiloxane network is substantially unstrained. That is, while parturbed polymer chains or networks can be obtained by swelling or the application of other shear stress to the polymer after formation, such as dissolving the formed polymer in a solvent, the resulting swollen network will be strained and thus undergo less deformation and have greater retractive stress than a relaxed network such as formed according to the present invention.

Surprisingly, it has been discovered according to the present invention that the hydrosilylation reaction

between the silicon hydride and vinyIslioxane groups occurs even where the reactive solvent (monomer) contains vinyl or vinylidene groups, and even where the concentration of vinyl groups on the vinylislioxane is extremely low. This was quite unexpected since the hydrosilylation reaction has been reported to take place between SiH and vinyl or vinylidene groups such as in methylimethacytels and acrylate monomers (F. P. MacKay et al., J. Am. Chem. Soc. 78:2784 (1975)) and in styrene and alpha-methylstyrene (J. W. Ryan et al., J. Org. Chem. 24:2052 (1959)) using a typical platinum-based hydrosilylation catalyst for this reaction at ambient or elevated temperatures. Accordingly, the present invention may be carried out with substantially no cross-linking between polyorgenosiloxane and the monomer for the second and any additional polymers present in the composition.

The preparation of polyorgenositoxianes by hydrosilylation (sometimes reterred to as two-part vulcanizing silicones) is well known in the art, and the formation of such systems is described, for example, in Arides patient 4,500,668. In general, the two-part system generally contains one polymeric silicone component having silicon hydride (SiH) groups, while the other polymeric component contains unsaturated groups, preferably viruly groups. Reaction of the silicon hydride groups with the unsaturated groups results in cross-linking of the components. Both the hydride and unsaturated groups can be part of the same polymeric silicone, in which case the silicone cross-links with itself.

Generally, such reactions are carried out with a catalyst, preferably a plathnum complex. Virtually, any of the conventional silicone components used in forming polyogenacilicances by hydrostilylation may be used in the present invention. Preferred organisationshares for use in the present invention include polydimethyl-so siloxanes, polymethyl-phryslicxanes, and various copolymethyr-methylationshares, polymethyl-phryslicxanes, and various coopolymers threach. More particularly, vinyl-terminated polyslicxanes are preferably reacted with methylhydrosiloxanes containing at least two hydrides are molecule.

The monomers which serve as the dispersion medium for the siloxane components and which react to form the second polymer or any additional polymer may be selected from any of a wide range of a monomers which will not inhibit the hydrosilylation reaction. Such monomers include those which undergo free radical, anionic, cationic or organometallic polymerization. Particularly preferred are monomers which undergo free radical addition polymerization, particularly vinity and vinyridene monomers.

While it will be understood from the foregoing that the invention is not limited to any particular second or subsequent polymer. The invention will be described heresider, with particular reference to vinyl and so vinylidene polymers. Particularly preferred monomers for use in the present invention include the acrylates, methacrylates, styrenes including e-methylstyrenes and alkylstyrenes, whyl ethers, acrylamides such as N-alkylacry-to methacrylamics, whyle sters, whyle prefelches and whyl pyridines. Still more preferably, the monomer is a homogeneous whyl monomer system which allows the preparation of optically clear compositions according to the present invention.

The hydrositylation reaction may be represented by the following equation which shows a single cross-

wherein R' through R<sup>5</sup> can independently be any of a wide variety of organo groups which are used in siloxane polymers, including saturated or alkyl or aryl, with partial or complete halogen substitution, evanositivil or evanoprovil, amino-substituted arryl or alkyl groups, and the like; and n and m indicated

degree of polymerization which may vary from 1 to about 100,000,000.

As indicated above, the reaction is preferably catalyzed by a platinum catalyst such as those described in U.S. patents 3.159,601; 3.159,662; 3.419,593; 3.715,334; 3.775,452 and 3.814,730. A particularly suitable catalyst is a complex of hexachloroplatinic acid with tetramethyldivinyldisiloxane, which may be present in a namount of less than about 1 percent by weight of the total composition.

As the hydrosilyation reaction proceeds, a significant increase in viscosity is observed in the organosiloxane/monomer solution or dispersion. The rate of viscosity increase varies depending upon a number of factors, including: (1) the polysiloxane components present, (2) the vinylsiloxane and silicon hydride concentrations, (3) the reactor conditions of temperature/pressure and degree of mixing, and (4) the of degree of cross-links formed by the hydrosilyation reaction.

The viscous solution or dispersion eventually forms a got, and the degree of gelation may be terminated at any given viscosity by adding someon sense, thinkblor or a retardant for the hybriardistation renational of desired, the get at this point may be cast into a mold or formed into a coating or film prior to the next reartion sten.

The next process step consists of converting the organic monomer or monomers into polymeric material, preferably by means of a free radical addition polymerization machanism. This polymerization may be carried out at ambient or elevated temperatures with a free radical initiators such as an azo or peroxide initiator. Unless it is desired to also cross-link the second polymer at this point, it is preferred to use azo or other low temperature initiators in order to avoid cross-linkino.

and Many suitable free radical initiators are known in the art and commercially available, including, for example, butyli peroxypivaleals, a low temperature peroxide catalyst sold under the trademark Lupersol-11 (a 75 percent catalyst solution in mineral spirits), 1,1-Bist/butyl peroxy) 3.3.5-t/methylcyclohexarie sold under the trademark Trigonov-29 (75 percent active), 2,2-Acobis (2.4 dimethylvaleronitely) and 22 Acobis (isobutyronitrile) sold by E.I. DuPont de Nemours and Company under the trademarks Vasc-62 and Vazo-84, respectively. Alternatively, the free radical polymerization may be initiated by ultraviolet light using a photosensitizer such as isobutylbenzoin ether, available from Sloutfer Chemical Company under the trademark Vicura-10.

For a most efficient polymerization, it is usually preserved to use at least two different initiators, which are activated at different temperatures. That is, after activating and using up a first initiator at a first temperature, the polymerization may be accelerated by raising the temperature to activate a second initiator. When the second initiator is virtually used up, the temperature is usually raised again to consume all of the initiators and complete the polymerization.

The above-described two step casting process is particularly suited to the molding of complex, fiexible medical parts as well as ultraviolet cured coatings. The pre-gelation step allows good control of the 1st relocity of coatings which is an extremely important factor in high speed coating processes. Also, as noted above, the free radical polymerization of the monomer may be completed, for example, inside a closed mold to produce rigid or flexible molded articles.

Although the above process has been described as a sequential polymerization, it will be understood the hydrostlylation reaction and monomer polymerization could be carried out simultaneously, for example, by raising the temperature of the hydrosilylation reaction to the point where at least one of the free radical initiators is activated to effect polymerization of the monomer. However, the two-step sequential process is preferred to maintain process control and evoid undestrable cross-inking polymers.

Without cross-linking of the second polymer, the above process produces single or mono-penetraling networks (MPN) of polysiloxane, if desired, the second polymer may be cross-linked by the inclusion of a cross-linker, such as a multi-functional vinyl or vinylidone molety, in the reaction mixture. Suitable cross-linking agents for vinyl and other polymers are well known in the art, including, for example, divinybenzene, dimethacrylate esters, and methacrylic acid, which are particularly suitable for cross-linking methacrylates or styrenes during polymerization.

Coss-linking of the second polymer results in the production of a second polymer network which interpenetrates the polystioxane network, thereby forming duel-penetrating networks (DPN). If a third or subsequent polymer is included in the reaction mixture and cross-finked to form separate polymer networks, the composition may be a triple penetrating network (TPN) or more. In general, the invention relates to preparing poly-penetrating networks (PPN) wherein at least one of the networks is a polyogenacilloxane network formed in a multi-component network system. The other polymers may be thermoplastic or selastomeric, although the curing will usually result in the formation of a thermosetting composition.

In addition to the above-described modified-bulk polymerization method, the present invention may be carried out with a number of different variations which are described below. One variation which is particularly suitable for the preparation of ele-locatings involves carrying out the hydrosilytation step in a

homogeneous mixture of vinyl monomer, an organo multi-functional cross-linker and an inert solvent, preferably an aromatic solvent such as toluene, hexane, heptane, cyclohexane, a parafinic solvent such anaphto, or any solvent which would not interfere with platinum catalyst. The cross-linking of the siloxane gell (hydrosilytation) takes place at ambient temperatures, followed by free radical polymerization of the monomer at ambient or elevated temperatures.

By adjusting the type or amount of whyl cross-linker, a flowable coating with a controllable rheelogy can be produced. Further, prior to or during the free radical polymerization, an ionic and/or polar winyl monomer may be incorporated into the reaction mixture to produce organositoxane coatings with excellent adhesion to concrete, glass, rubber, metal and other surfaces. Film coatings from these organositoxane interto penetrating networks show inusually high elasticity and excellent recovery, making these coatings very suitable for cating onto other elastomers and flexible tubino.

The inherent mechanical, gas permeability, optical and other properties of these films, membranes and coatings prepared by casting flowable interpenetrating polymer solutions of the present invention may be adjusted by varying the composition of the organic monomer phase. For example, by introducing 16 hydrophilic monomers into the reaction mixture, hydrophilic organisations are particularly suitable for medical applications. Similarly, by adjusting the preferctive index of the vinyl monomers and the polyslioxanes used, as well as by controlling the cross-link density, clear or opaque coatings may be obtained.

In another embodiment of the invention, a suspension polymerization process may be used. For example, the hydrosilyation reaction may be carried, out in a primary reaction in the presence of vinyl monomer and free radical initiator. After completion of the gelation of the polysiloxane elastomer, the solution is dispersed in a second reactor in the presence of a colloid, or preferably at a temperature where the half-life of the free radical initiators used is about 1-2 hours. A fine dispersion of particles is produced, and the polymerization of the monomer is carried out with one or more initiators. After polymerizing the vinyl ze monomer, small amounts of talc or silica or other blocking apents are added to the reaction before the situry is allowed to cool down. This produces rubbery beads which should be air dried before being molded or extruded into a final rigid or elastometric product.

An important advantage of the compositions of the present invention is that all of the ingredients, with the possible exception of the platiam catalyst, may be mixed together and stored or transported for so considerable periods of time until ready for reaction and transformation into the interponentating polymer networks of the invention. For example, a premixture containing the vinysibiliscane and sition, hydride components, the organic monomer, the azo or peroxide initiators and the multi-functional vinyl cross-linkers may be prepared and stored until reaction is desired. The premix may then the treated with the platinum catalyst and heat at the desired level or levels to effectuate the hydrosilytation, polymentzation and/or cross-si linking, either sequentially or simultaneously. As desired.

The compositions of the present invention yield final products with improved dimensional stability and elastic behavior over the standard filled silicone rubbers. Further, since the vinyl polymerization step can be completed in a matter of a few hours instead of many hours of oven "post-curing," a significant improvement in process and handling can also be achieved. Controlling the modulus of the second penetrating network via different cross-linkers, the Shore A hardness and stiffness can be varied with a Shore A or Shore 00 from 0 to 100. The modulus and hardness may also be adjusted by selecting the proper vinyl or vinylidene monomer mixtures. Still further, the amount of the cross-linker used during the second stage can provide opaque to clear castings and moldings.

Improved biocompatibility of the compositions of the invention over conventional silicon materials can be achieved by incorporating amino acids and polar groups into the second polymer network. Also, as anoted above, the present invention provides an improved route for producing silicone membrane materials with good permeability, which is externely important in gas separation and biomedical application. Combined with the ability to produce optically clear films and coatings, the permeable membrane materials are particularly useful, for example, in the manufacture of contact lenses.

The present invention will now be illustrated in more detail with reference to the following specific, nonlimiting examples. In each of the examples, the catalyst used was a toluene solution of hexaphorpalaritic actiodivinylitetramethy/distloxane complex containing 0.35 weight percent platinum (commercially available from Pétrarch Systems Inc. under the designation PC072). All parts are parts by weight, but percentages in the siloxane compounds are mole percents.

This example covers bulk polymerization of the compositions outlined in Table 1. The general polymerization procedure was carried out in three steps. After a homogeneous solution was prepared the platnum catalyst was added, allowing the hydrosilylation reaction to proceed for three hours at ambient temperatures under nitrogen until the polyorgenosiloxane network was formed. The second stage was so carried out in horizontally placed trays producing sheets for determining ASTM physical properties. The bulk polymerization experiments in Table 1 produced only solid polymer DPM systems if the platimum catalyst was added, demonstrating that the hydrosilylation reaction was critical in order to obtain a useful product. Eliminating the platimum catalyst and thus preventing the hydrosilylation reaction from taking place, we observed only oily, heterogeneous, greasy fluid products after completing the free radical reaction (Table 1 and 2, Flurs 5 and 6). The semi-interperetating networks (MPN) had intipher elongation than the full dual-penetrating network systems (DPN). The tensis strength and hardness increased by going from an MPN to a full DPN. These examples demonstrate a novel route for preparing siloxane elastomers without going through expensive compounding and crossilishing operations.

### Example 2

20 This example covers preparation of semi-penetrating networks according to the procedure of Example 1 where the primary network its prepared from a sliowan entwork (Table 3). When the second polymer was prepared from a linear styrenic polymer, the properties of these elastomers verified that a highly elastic material may be prepared. The t-bulyt-styrene processed system (SPR) showed excellent optical properties. The properties in Table 4 show unexpected elongation and high resilience over standard silicone subtractival.

### Example 3

This example describes dual penetrating networks (PPN) of PDMS/PS, prepared according to Example 1 with the compositions and conditions of Table 5. Physical data tabulated in Table 5 verified that hardness and tenalle atrength were significantly improved by diviyotherazene. The tent-butylstyrene polymer was found to produce unexpectedly good compatibility with the polydimethylstions ne nervoir. The crosslinking via hydrosilyation was found to be critical for producing a useful product since the vinyl terminated siloxane and the silicone hydrides do not participate in a free radical reaction at the polymerization conditions call. Eliminating the platinum catalyst produced liquid heterogeneous viscous fluids with separate domains of some solid oplystyrenic againementae (Run 14).

### Example 4

This example demonstrates how optically clear, semi-penetrating networks of polydimethylsiloxanes, objymethyleheylsiloxane copplymers and polymethylarysilect esters may be prepared via the present invention. Using a copolymer of dimethyldichlorosilane and diphenyldichlorosilane or copolymer prepared from octamethylcyclothersiloxane and calophenylcyclothersiloxane or polymethylphenylsiloxane, having at least two vinyl groups per molecule, pre-dissolved in methacrylatio or acrylate esters together with a nutlithyldride sloxane, clear water-white moldable polymers were prepared after carrying out the golation at so ambient temperature for two hours, followed by three hours at 75°C and two hours at 115°C. A typical example is given in Table 17 (run 17), where a copolymer of dimethyl(175%)-dimethyl(185%)-sloxane (170%) termination having a refractive index of 1.40 was cross-inked with methyllydro(30%)-dimethylsioxane(170%), products are suitable for contact lens materials, medical tubing and other optically clear products, prepared by polymerization casting or extrusion-molding technology.

# Example 5

This example demonstrates how the invention produces obtically clear penetrating polymeric systems from incomposible, opaque systems such as polydimethylsioxane/polymethacy/late seters, where the stractive indices are 1.40 and 1.48 and the cohesive energy densities are about 15(Jcm²)<sup>1/20</sup> and 18(Jcm²)<sup>1/21</sup> for the corresponding polymer systems. Using the compositions of Table 8, we discovered that completely clear material can be prepared from opaque incompatible systems as described above. The physical properties presented in Table 8 demonstrate that the crosslinking density of the second network can be adjusted to produce clear materials. Properties such as break stress, handriess, tear strength and elastic recovery are also improved over silicone elastomers using the dual network approach. This invention demonstrates how low-cost polydimethysicoanepolymethacytate esters can produce clear, castable or moldable optical materials, ranging from soft to rigid materials which may be molded or shaped into any useful product.

### Example 6

This example demonstrates how the present invention can be extended to cover forometic interpenentaring network systems. Runs 23 and 24 shown in Table 10 cover dual penetrating networks of polydimethylsiloxane/polystyrene systems where the polystyrene network is produced by covalent (Run 23) or incomeric. (Run 24) bonding compared to a non-cross-linked system (Run 22). The use of inomeric bonding rather than covalent bonding allows production of more llowable, dual-network polymeric material and had a Shore A hardness of 84 (Run 24). The fact that platinum catalysts allow the hydrosilylation reactions for proceed even in the presence of ionic vinyl monomers was also quite unexpected, especially since platinum catalysts for the hydrosilylation reactions are known to be deactivated by strong scids. The internal hydrogen bonding between two acrypic acid monomers may have overcome this problem, thus allowing extension of the invention to dual innomeric interpenetrating networks as demonstrated in this example.

# Example 7

This example illustrates the preparation of dual interpenetrating networks of polydimethylsiloxane and polyvinylisobutyl ethers, where the second network poly(vinylisobutyl) ether consists of a semi-crystalline thermoplastic network, while the primary network (the PDMS network) is a permanent network. The following polymerization was carried out in bulk under nitrogen on a 6 x 6 in2 Tellon coated tray. Twenty grams of a vinyl-terminated polydimethylsiloxane, having a kinematic viscosity of 165,000 ctsk, were 40 dissolved in twenty grams of vinylisobutyl ether, producing a homogeneous solution. To this solution was added 2.5 grams of a methylhydro(30%)-dimethylsiloxane(70%) copolymer having a kinematic viscosity of 1000 ctsk. After cooling this solution down to 0°C under nitrogen, 0.090 grams of the platinum catalyst (0.35 wt.%) and 0.040 grams of borontrifluoroethyl ether complex were quickly mixed, and the mixture was allowed to polymerize under nitrogen inside a Teflon coated tray, stored on ice. After four hours we 45 observed that a tough, rubbery, optically clear 72 mils thick sheet with a Shore A of 43 had been produced. This improved toughness resulted from the polydimethylsiloxane network produced via the hydrosilylation reaction and the semi-crystalline network structure present in isotactic poly(isobutyl)vinyl ether having a crystalline melting point of about 150°C. The dual network produced above can be molded or extruded above the melting point mentioned, having improved hydrophilic, permeability, adhesion and mechanical 50 properties compared to polydimethylsiloxane elastomers.

### Example 8

This example describes the preparation of interpenetrating networks in the form of flowable "gel coatings" by controlling parameters such as molecular weight and cross-link density in each of the separate networks

To a stirred reactor was charged fifty parts of a viny-terminated siloxane having a kinematic viscosity of \$5,000 ctst, three parts of a (30%)methythytic-70%/sidemthytischerace pooplymer with a viscosity of 300 cps., forty-five parts of 2-ethythexylmethacrylate and fifteen parts of isobutylmethacrylate added under dry nitrogen. The hydrositylation reaction or the polysiloxane gelation reaction was allowed to proceed for two hours at ambient temperature before 0.080 parts of Vazo-92 and Vazo-94 were added. The reactor temperature was increased to 65°C, 80°C and 95°C in steps of two and one-half hours at each temperature with one-half hour heal-up time between each step. A free flowing figuid with kinematic viscosity of 87 centistokes at a shear rate of 16.8 reciprocal seconds at 23°C was produced. The physical properties of the coating were setted after casting a 10 mill film from the "glic cotating" onto a Teflon surface. The physical properties of the coating were setted after casting a 10 mill film from the "glic cotating" onto a Endon surface. This physical properties of the on dual-interpretating networks. This get coating is a stable, homogeneous, flowable poli, Addition of toluene as a diluent results in dual phases, one phase containing the Interpenerating network phase with a clear toluene phase above.

### Example 9

This example demonstrates the preparation of hydrophilic coatings by incorporating water soluble or 20 Ionic monomers during the polymerization step. To a reactor was added twenty-five parts of vinyl terminated polydimethylsiloxane having a viscosity of 16.500 centistokes, three parts of a 25 centistoke (30%)methylhydro-(70%)dimethylsiloxane copolymer, twenty-five parts...of 2-ethylhexyl-methacrylate, two parts of N-vinyl pyrrolidone, 0.32 parts of diethylene glycolmethacrylate and one hundred parts of toluene. Furthermore, 0.1 parts of a 0.35 weight % platinum catalyst together with 0.050 parts of Vazo-52 and Vazo-25 64 were added. After completing the silicone crosslinking reaction (i.e. the hydrosilylation reaction) the free radical reaction was carried out at 65C, 80°C and 95°C over a time period of two and one-half hours at each temperature, with a heat-up time of one-half hour. The final product was a white stable fluid with a solids content of 34% having an apparent viscosity of 281 centipoise at 23°C measured on a Brookfield Viscometer at a shear rate of 168sec-1. Physical properties were measured on a 10 mils thick film cast from the reactor product. The completely elastic film had a break stress of 162 psi, elongation at break of 617%, set at break of 0%, Shore A Hardness of 26. The casting showed excellent adhesion to glass, metal and flexible plastics as well as human skin. Immersing 10 mil tensile bars of pure silicone gel coating (control) and our hydrophilic network coating in distilled water for thirty minutes followed by thirty minutes air drying verified a water uptake of 4.69 weight % for our hydrophilic coatings vs. 0.07 weight percent for the control, thus confirming the improved hydrophilic nature of the dual interpenetrating network coatings according to the present invention.

# Example 10

This example describes the preparation of release coatings based on dual penetrating networks of polyorganositoxera end boymethyl-imbercytate setters for high speed coating applications. The following solution polymerization was carried out. Viryl terminated polydimenty/siloxane, fifty parts, having a bulk viscosity of 155,000 centrolitoses at 23° C together with three parts (20%-inethylhydror-(70%-johrentylsiloxane opportunity) of 155,000 centrolitoses and solution viscosity of 25:300 centrolitoses, were dissolved in 250 parts toluenes. Goldenot by the addition of 37.5 parts of 2-ethylhexylinethacrylate, 0.325 parts of divhylbenzene and 12.5 parts of isobultyinethacrylate. To this solution were added 0.150 parts of a 0.55 weight % platinum catalyst and 0.050 parts of 2.2 -Azobis(sobulyrontrile). Following three hours of hydrositylation at 25° C, the fire artical polymerization was carried out at 80° C, 80° C and 90° C for two hours at each step under nitrogen. Final product had a solids content of 28° weight percent with a kinematic viscosity of 280 centistokes. A 0.5 mil coating on alaminum substate produced a release value of 82 gramsin. For a styrene-butadiene rubber adhesive-coated label tested with a tweet inch per minute strain rate with a 180° peal angle, using a one-inch wide coated label. Adjusting the acrylic monomer ratio (tacky acrylics below glass transition temperature(T<sub>2</sub>) versus hard acrylics above T<sub>2</sub>) in this example, coatings were prepared argning from low release to adhesive properties.

### Example 11

This example illustrates the preparation of a concrete sealing coating based on our invention. To a 5 reactor was charged fifty parts of a vinyl terminated siloxane having a viscosity of about 65,000 centistokes. six parts of a (30%)methylhydro-(70%)dimethylpolysiloxane with a viscosity of 25-30 centistokes, twentyfive parts of 2-ethylhexylmethacrylate, twenty-five parts of isobutylmethacrylate, toluene 250 parts. After 0.180 parts of the platinum catalyst (0.35 weight %) was added the gelation reaction was allowed to proceed for two hours at 23°C under nitrogen, before 0.325 parts of divinylbenzene and 0.50 parts of Vazo-10 52 and Vazo-64 were added and the polymerization completed as in the Example 9. The final solution had a solids content of 27.3 weight % and a solution viscosity of 48 centistokes. These semi-translucent coatings were applied to dried 4" x 4" x 4" concrete blocks and Immersed in deignized water with a control block in a separate bath. After twenty-four hours of immersion the control picked up 3.8 weight % water vs. 0.6 weight percent for the coated material. After thirty days immersed in water the uncoated concrete block 15 showed significant signs of erosion, while the coated block remained unchanged. The excellent performance of the concrete coating was also verified by no changes in the pH of 7, while the immersion bath for the control had changed to a pH of 8.2 and to a yellow color with concrete debris present on the bottom. The sealant showed one-hundred percent performance after sixty days while the control block was now developing severe pitting.

### Example 12

This example demonstrates the preparation of dip-coating, semi-permeable and other highly, elastic membranes utilizing the present invention with dual interpenetating polydimethysticioxen and addition polymers. To a reactor was charged fifty parts of dimethysilicioxen vinyl terminated, four parts of (30%)-methyshydro-(70%)climethyl, twenty parts of 2-ethylhexylmethacylate, thirty parts of isobudy/methacylate and two-hundred parts of tolene. The hydrostilyation reaction was activated by 0-150 parts of a 0.3 weight percent platinum catalyst and completed over the next two hours at 23°C. Thereafter, 0.55 parts of divilybenzene together with 0.5 parts of Vacc>25 and Vacc>64 were added, before the polymerization was completed at 65°C, 80°C and 95°C for two and one-half hours at each step. The final product had a viscosity of 550 certipoles, Coatings applied horizontally or vertically to Telfon surfaces produced highly homogeneous, highly elastic membranes, with excellent elastic recovery. Coatings were compared to pure found to be inferior in mechanical and rheological properties compared to the interpenetrating network coatines of the present invention.

### Example 13

This example Illustrates the preparation of pressure sensitive adhesives based on dual penetrating networks of the present invention. To a stirred nearctor was charged lwenty-five parts vinyl terminated polydimethylsiloxane with a kinematic viscosity of 65,000 centistokes, three parts of (30%-imethylydroc/(70%-idimethylsiloxane copolymer of 25-30 centistokes, twenty-five parts of 2-ethyfixey/methacrylate and two-hundred parts of tolivane. After adding 0.08 parts of 0.30 weight % platimum catalyst, the gelation step was extended over two hours at ambient temperature. Next, 0.50 parts of Vazo-52 and Vazo-44 together with 0.65 parts of divinybrance were added before the polymerization was carried out at 55-00 °C, 75-08 °C C and 95 °C for two and one-half hours at each step producing a semi-translucent fluid with a viscosity of seventy-eight centistokes. Autimium foil, 6 mils thick, was coated with a one mil thick lim of final product before being rolled together with a 5 pound weight face to faze. A lap peel test was run according to ASTM-D-1000 showing a peel adhesion of 612 g/cm, which remained constant with time. This demonstrates that a pressure sensitive adhesive can be prepared directly from coatings of the invention, 59 without requiring the time sensitive curing process normally required for pure silicone-based pressure sensitive adhesives.

### Example 14

This example illustrates the preparation of coatings or membranes by a continuous process using ultra violet radiation polymerization on a continuous belt or in a continuous coating, casting or calendering operation. To demonstrate this process, we used a Fusion System's, Model F480, equipped with a continuous belt moving at a constant rate, having a U.V. lamp (D bulb) mounted above. A homogeneous polydimethylsikoane gel was prepared in a methacrylate sette by carrying out the pre-gelation step in a tel through the following procedure: fifty parts of vinyl terminated polydimethylsiloxane having a tokinematic viscosity of 65.000 centistokes, our parts of (30%)methylsiloxane copolyme, fifty parts of isobulytimethoxide, 0.65 parts of diethyleneglycolmethacrylate and two parts Vicure-10 were mixed until a clear homogeneous solution was prepared. Next, 0.180 parts of a 0.35 weight percent platinum catalyst were quickly mixed in, and the solution cast onto a flat surface, allowing the hydrositylation reaction to proceed until a non-flowable gel was produced. The gel was passed under the U.V. lamp at 16 a rate of 11 firthmia lallowing a liexible membrane of 5 mit. Inthicness to be produced. This highly elastic membrane of dual interpenetrating network polymer had significantly improved mechanical properties compared to bulk hydrosilystom which resulted in an extremely soft and fulled polysiloxane compositions.

It will be recognized by those skilled in the art, that changes may be made to the above-described embodiment of the invention without departing from the broad inventive concepts thereof. It is understood, therefore, that this invention is not limited to the particular embodiment disclosed, but it is intended to cover all modifications which are within the scope and spirit of the invention as defined by the appended claims.

Table 1

Proparation of Memogenetrating (RPM) and Dual Penatrating (RPM)

Metworts of Polygiloxane and Polygilyronic Materials

Chemicals	Run-1	Run-2	Run-3	Run-4	Run-5	Run-6
Type of 1PN -	NDW	NAG	мьи	NAO	Control	Sontrol
Methylhydro(J03)- dimethyl siloxane(703)- copolymer, visc. 25 ctsk., perts		٠.	īv	2	, <b>v</b>	rs.
Polydimethylsiloxane, vinyl terminated, visc., 1000 ctsk., parts	. 6	40	40	40	40	0
tert-butylstyreno, parts	40	0		i		٠.
styrone, parts		•	40	40	40	40
dlivinylbenzene, parts		-		-		-
Supersol-11, parts	0.075	0.075	0.075	0.075	0,075	0.075
Trigonox-29, parts	0.015	0.015	0.015	0.015	0.015	0.015
Pt-catalyst (0.35 wt.8), parts	0.060	0.060	0.060	0.060		

oily

elastomeric

elastomeric

elastomeric

elastomeric

Product Description

Table 2

Physical Properties of Monopenetrating (MPN) and Dual Penetrating Network (DPN) Systems of Polysiloxane-Polystyrene Materials

	Properties	Break* strength psi	Elongation at break 3	Regilience Hardness ASTM-D- Shore - 2632 A OO	Hardness Shore - A OC	00 00	Type of (IPN) Inter- penetrating Network Systems	
Run No. Run-1		82	692	48	6	5.4	MPN	
Run-2		312	220	35	. 09	68	DPN	
Run-3		137	738	. 95	7	39	MPN	
Run-4		255	. 539	. 43	17	88	NAO	
Run-5		fluids			1		,	
Run-6		fluids		*	•	1 8	÷	

\* ASTM-D-638 Crosshead speed 2"/min.

Table 3

Preparation of Monopenetrating Network Materials (MPN) of Polydimethylsiloxane and Organo-vinylpolymers

siloxane(?0%)copolymer, 25 ctak, parts 5 5 5  Polydimethylsiloxane, vinyl terminated, 1000 ctsk., parts 40 40 40 tsrt-butylstyrene, parts 40	Run No.	Run-7	Run-8	Run-9	
Polydimethylsiloxane, vinyl terminated, 1000 ctsk., parts 40 40 40 tsrt-butylstyrene, parts 40 Styrene, parts - 40 - 1so-butylmethacrylate, parts - 40 Lupersol-11, parts 0.075 0.075 0.075 Trigonox-29, parts 9.045 0.045 0.045	Chemicals .		*		
### ##################################			<del></del>		
25 ctsk., parts 5 5 5 Polydimethylsiloxane, vinyl terminated, 1000 ctsk., parts 40 40 40  tert-butylstyrene, parts 40 Styrene, parts - 40 iso-butylmethacrylate, parts - 40 Lupersol-11, parts 0.075 0.075 0.075 Trigonox-29, parts 9.045 0.045 0.045	Methylhydro(30%)-dimethyl				
Polydimethylsiloxane, vinyl terminatud, 1000 ctsk., parts 40 40 40 40 40 40 40 40 40 40 40 40 40	siloxane(70%)copolymer,				
vinýl termineted,       1000 ctsk., parts     40       40     40       40     -       -     -       Styrene, parts     -       iso-butylmethacrylate, parts     -       Lupersol-11, parts     0.075     0.075       Trigonox-29, parts     9.045     0.045	25 ctsk., parts	. 5	5 .	5	
1000 ctsk., parts 40 40 40 test-butylstyrene, parts 40 Styrene, parts - 40 - Iso-butylmethacrylate, parts 40 Lupersol-11, parts 0.075 0.075 Trigonox-29, parts 9.045 0.045 0.045					
tert-butylstyrene, parts 40 Styrene, parts - 40 iso-butylmethacrylate, parts 40	vinyl terminated,				
Styrene, parts - 40 - iso-butylmethacrylate, parts - 40  Lupersol-11, parts 0.075 0.075 0.075  Trigonox-29, parts 9.045 0.045	1000 ctsk., parts	40	40	40	
Styrene, parts - 40 - iso-butylmethacrylate, parts - 40  Lupersol-11, parts 0.075 0.075 0.075  Trigonox-29, parts 9.045 0.045					
iso-butyImethacrylate, parts 40  Lupersol-ll, parts 0.075 0.075  Trigonox-29, parts 9.045 0.045	tert-butylstyrene, parts	40		-	
iso-butyImethacrylate, parts 40  Lupersol-ll, parts 0.075 0.075  Trigonox-29, parts 9.045 0.045					
Eupersol-11, parts 0.075 0.075 0.075 Trigonox-29, parts 9.045 9.045 0.045	Styrene, parts	-	40	-	
Eupersol-11, parts 0.075 0.075 0.075 Trigonox-29, parts 9.045 9.045 0.045	V •				
Lupersol-11, parts         0.075         0.075         0.075           Triconox-29, parts         9.045         9.045         0.045	iso-butylmethacrylate, part	s -	8 =	40	
Trigonox-29, parts 0.045 0.045 0.045		-			
Trigonox-29, parts 0.045 0.045 0.045	funereol=11. parte	0.075	0.025	0.025	
	supersor ir, pures	0.075	0.075	0.075	
	m : -: «»				
Pt-catalvet (0.35 wt. %)	irigonox-29, pares	5.045	0.045	0.045	
parts 0.060 0.060 0.060		0.060	0.060	0.060	

Polymerization conditions similar to those described for Table 1.

Table 4

Physical Properties of Monopenetrating Network Materials (MPN)

# of Polydimethylsiloxane and Organo-vinyl Polymers

Hardness Tear** Resilience Molt*** Shore Die B ASTM Flow A 00: 1b/in D2632 g/10 min	34 78 159 36 1.74	6 50 225 28 1.12	16 50 302 -
Blong.* at break %	972	. 226	763
Break* Stress psi	305	392	513
Properties Break* Stross Psi Run No.	Run-7	Run-8	Run-9

\*ASTW-D-638 at 2 in/min.
\*\*ASTW-D-624 at 20in/min.

\*\*\*ASTM-D-1238 G conditions

Table 5

Proparation of Dual Peneirating Metworks of Polydimethylailoxane and Polystyrenics via Sequential Polymerization

Run No	Run-10	Run-11	Run-12	Run=13	Run-14	
Methylhydro(101)dimathyl- siloxane(701)copolymer, 25 ctsk., parts	. 5	ž	5		۰, ۰	
Polydimethylsiloxane, vinyl terminalod, 1000 clsk., parts	40	40	. 04	. 04	40	
Styrene, parts	40.	09	•	40	40	
tert-butylstyrene, parts	•	•	40		١,	
Divinylbenzenc, parts		9	4	,	4	
fupersol-11, parts	0.045	0.060	0.045	0.045	0.045	
Trigonox-29, parts	0.045	0.045	0.045	0.045	0.045	
Pt catalyst(0.35 wt.%), parts	0.060	0.060	0.060	. ,	٠,	

2 hrs. at 25% 3 hrs. at 75% 2 hrs. at 110

Polymerization conditions: (similar to description in Example 1)

Table 6

15

of Polydimethylsytrene and Polystyrenics with High Degree of Crosslinking Physical Properties of Dual Penetrating Network Materials (DPN)

Run No.	Properties	Tensile* Break strength psi	le* gth	Elongation at Break %	Tear** Die B ft lbs./in.	Shore A*** Hardness	Sett at Break %
Run-10		588		42	37	79	1.
Run-11		694		40	39.5	. 83	
Run-12		325		848	. 02	63	1.7
Run-13		Too f	luid	Too fluid and hetergeneous to test.	ous to tes	i.	
Run-14		Too f	luid	Too fluid and hetergeneous to test.	ous to tes		

\*ASTM-D-638 at 2 in/min \*\*ASTM-D-624 at 20 in/min \*\*\*ASTM-D-2240 - Durometer Hardness at 23<sup>O</sup>C

Table 7

Preparation and Properties of Monopenetrating Networks of 
Polysiloxanes and Polymethacrylate Esters (MPN Materials)

Run No.	Run-16	Run-17	
Formulation			
Methylhydro(30%)-dimethylsiloxane (70%)coplymer, 25 ctsk., parts	2.5	2.5	
Polydlmethylsiloxane, vinyl terminated, 1000 ctsk., parts	. 20	-	
Dimethyl(75%)-diphenyl(25%) siloxane copolymer, vinyl terminated, 1500 ctsk., parts	-	20	
Isobutylmethacrylate, parts	20	20	
Lupersol-11, parts	0.060	0.060	
Lupersol-256, parts	0.045	0.045	
Pt-catalyst (0.35 wt.%)parts	0.030	0.030	
Properties	•		
Hardness Shore A	67	82	
Resilience ASTM-D-2632	55 -	43	
Optical Clarity	turbid	, clear	

Run-21

Run-20

Run-19

. Run No. - Run-18

Chrmicals				
Polydimethylsi loxant vinyl cerminited, 1000 cesk., parts	20	95,	. 95	9.0
Methylhydrollun)- dimethylsilvxanm{748}- cupolymer, 25 ctsk., parts	v		VE	
2-ethylbexylmuth- ucrylate, parts	20	20	. 20	. 05
Divinylhenzene, parts	,	-	2	3.5
Lupersol-11, parts	0.120	0.120	0.120	0.120
Trigonox-29, parts	0.060	0.060	090.0	0.060
Pt-catalyst(0.35 wt.2), parts	0.180	0.180	0.180	0.180

Polymerization Conditions: (similar to description for Table 1)

2 hours at 25°C 3 hours at 75°C 2 hours at 115°C

Physical Properties of Clear, Dual Penetrating Network of Polydimethylsiloxane/Polymethaciylate Haterials

Properties	Break (1) Stress psi	Flongation (1) Set (4) at Break at Break	Set (4) at Break 3	Tear (2) Die B 16/in	Shore A	£ 20	Shore A <sup>(3)</sup> Reliactive Optical Hardness Index Clarity A 00 n <sub>D</sub> (23°C)	Optical Clarity
Run No.								
Run-18	01	1700		17 0			white	anhedo .
Run-19	146	115	1.4	69	78 6	00	60 1.4290	clear
Run-20	241	420	1.6	. 16	36	80	1.4356	clear
Run'-21	473	293	ji	201	42 .8	. 85	1.4469	clear
			(1) ASTM-D	(1) ASTM-11-638, Crosshead speed 2"/min.	speed 2"	nim/		

ASTM-D-624, Crosshead speed 20"/min. (5) ASTM-D-2240, Durometer hardness at 2100 Ē

ASTM D-638 at 2"/min. Ξ

Table 10

# Penetration of PDMS/polystyrene Dual Penetrating Networks based on Ionomeric and Covalent Crosslinks

W	Run 22	Run 23	Run 24
Chemicals			
Polydimethylsiloxane			
vinyl terminated, 1000 ctsk., parts	20.		. 20
1000 Ctski, pares	20,		. 20
Methylhydro(30%)-			
dimethylsiloxane(70%)-		•	
copolymer, 25 ctsk., parts	5	5	5
	-		
Styrene, parts	20	20	2-0
Divinylbenzene, parts	-	1.5	- ·
W. Ph			. 1
Methacrylic acid, parts	-	<u>-</u>	
Lupersol-11, parts	0.045	0.045	0.045
napatata in parat		· ·	
Trigonox-29, parts	0.015	0.015	0.015
Platinum catalyst			
(0.35wt.%), parts	0.045	0.045	0.045

Polymerization conditions under nitrogen: (similar to those described in Example 1)

<sup>2</sup> hours at 25°C 3 hours at 75°C 2 hours at 105°C

### Table 11

Physical Properties of I Network Coatings o Polyorganosiloxane a Addition Polymer Netw	f nd
Tensile Properties *	
Break stress, psi Elongation at break % Set at break % Shore Hardness A**	125 700 0 28
Rheological Properties *** wt% in toluene)	(27
Shear rate, sec-1 Apparent viscosity, cps.	168 663

- \* ASTM-D-638 Crosshead 2"/min.
- \* ASTM-D-2240, Durometer hardness at 23
- \*\* Brookfield Digital Viscometer at 23.

### Claims

- A composition comprising a relaxed, perturbed, cross-linked polyorganosiloxane network dispersed in a second organic polymer.
- A composition according to Claim 1 wherein said second polymer is not significantly cross-linked, whereby said polygraphosiloxane forms a mono-penetrating network.
- 3. A composition according to Claim 1 wherein said second polymer is substantially cross-linked to itself but not significantly cross-linked to said polyorganosiloxane, whereby said polyorganosiloxane and said second polymer form a dual-penetrating polymer network.
- 4. A composition according to Claim 3 comprising at least a third polymer which is substantially cross-linked to itself but not significantly cross-linked to said other polymers, whereby said polymers form a polypenetrating polymer network.
- 5. A composition according to Claim 1 wherein said second polymer is an addition polymer.
- A composition according to Claim 5 will be acid addition polymer is selected from the group consisting of vinyl and vinylidene polymers.
  - A composition according to Claim 5 wherein said addition polymer is a glassy polymer having a glass transition temperature of about -120°C to 280°C.
- A composition according to Claim 5 wherein said addition polymer is a home- or copolymer having a melting point above 23\*C.
- A composition according to Claim 5 wherein said addition polymer is a thermoplastic ionometic polymer.
   A composition according to Claim 1 wherein said polyorganosiloxane is cross-linked by a
- hydrosilylation reaction.

  11. A composition according to Claim 1 wherein said polyorganosiloxane is present in an amount of
- 11. A composition according to Claim 1 wherein said polyorganosiloxane is present in an amount o about 0.01 to 99.99 weight percent.
  - 12. A composition according to Claim 1 which is optically clear.
  - 13. A composition according to Claim 1 which is homogeneous.
  - 14. A composition according to Claim 1 in the form of a gel coating.
- A composition according to Claim 1 in the form of a hydrophilic coating.
   A composition according to Claim 1 wherein said polyorganosiloxane is selected from the groups
- consisting of polydimethylsiloxanes, polydiphenylsiloxanes, polytrimethylsiloxanes, and copolymers thereof.

  17. A composition according to Claim 1 wherein said second polymer is selected from the group consisting of polyacrylates, polymethacrylates, styrenes, vinvloyrrolidones and vinvloyridines.

- 18. A method of making a cross-linked polyorganosiloxane network dispersed in a secohd organic polymer comprising dispersing a component having SIH groups and an organosiloxane having vinyl groups in a reactive liquid mortomer of said second polymer, cross-linking said component and said organosiloxane by hydrosilylation to form said network in said monomer, and simultaneously or subsequently polymerizing said monomer.
- 19. A method according to Claim 18 wherein said SiH groups and vinyl groups are on the same organosiloxane.
- 20. A method according to Claim 18 wherein said SiH groups are on a low molecular weight to polymeric component and said vinyl groups are on a separate organosiloxane.
- 21. A method according to Claim 18 wherein said component, organosiloxane and monomer are dispersed in a non-reactive organic medium.
  - 22. A method according to Claim 18 wherein said hydrositylation is catalyzed by a platinum catalyst.
    23. A method according to Claim 18 wherein the polymerization of said monomer is selected from the
- group consisting of free radical, anionic, cationic and organometallic polymerization.

  24. A method according to Claim 18 wherein said dispersion is substantially homogeneous.
  - 25. A method according to Claim 18 wherein said polymerized monomer is cross-linked to lorm a dual-penetrating network with said cross-linked polygraphosiloxane.
- 26. A method according to Claim 25 wherein at least a second monomer is polymenzed and cross-linked subsequent to said hydrositylation to form a poly-penetrating network.
- 27. A method according to Claim 25 wherein the cross-linking is carried out in a non-reactive solvent and the cross-linking density of each network is controlled to produce a smooth, flowing gel having a viscosity below about 10,000 centistokes at the boiling point of the solvent.
- dispersion droplets in non-reactive organic medium to form interpenetrating network beads, and recovering the network beads after addition of talc or silica to the medium.
  - 28. A method according to Claim 18 wherein said polyorganosiloxare network and said liquid monomer torm a gel which is then enuisitified in water with organosilicon block polymers, followed by emulsion polymerization of said monomer to produce a free-flowing aqueous coating of said network and said polymer.
- 30. A method according to Claim 18 wherein said network in said monomer is cast into a mold, and said polymerization is carried out after casting.
  - 31. A method according to Claim 18 wherein the monopenetrating network produced thereby is cast into a mold, and said polymerized monomer is cross-linked after casting.
- 32. A casting composition for coating and molding comprising a dispersion of a cross-linked polyorsg ganosiloxane in a non-silicone organic monomer, and an initiator dispersed in said monomer which will polymerize said monomer when activated.
  - 33. A casting composition according to Claim 32 wherein said initiator is activatable by heat.
- 34. A casting composition according to Claim 33 comprising at least two initiators activatable at different temperatures.
- 40 35. A casting composition according to Claim 32 wherein said initiator is selected from the group consisting of azo and peroxide compounds.
  - 36. A casting composition according to Claim 32 which also includes a non-reactive solvent for the components whereby the composition is in the form of a pourable gel for preparing films and coatings.
- 37. A casting composition according to Claim 32 which also includes a cross-linking agent for said so polymerizable monomer.
  - 38. A molding composition comprising dried beads for molding and extrusion, said beads comprising a dispersion of a relaxed, perturbed, and cross-linked polyorganosiloxane in a homogeneous, non-silicone organic polymer.